JAKAFI- ruxolitinib tablet Incyte Corporation
HIGHLIGHTS OF PRESCRIBI

BING INFORMATION

These highlights do not include all the information needed to use JAKAFI safely and effectively. See full prescribing information for JAKAFI.

JAKAFI [®] (ruxolitinib) tablets, for oral use Initial U.S. Approval: 2011	
RECENT MAJOR CHANGES	
Indications and Usage (1.3)	05/2019
Dosage and Administration (2.3)	05/2019
INDICATIONS AND USAGE	
Jakafi is a kinase inhibitor indicated for treatment of:	
• intermediate or high-risk myelofibrosis, including primary myelofibrosis, post-polycythemia vera myelofibrosis in adults. (1.1)	fibrosis and
• polycythemia vera in adults who have had an inadequate response to or are intolerant of hydroxyurea. (1.2)
• steroid-refractory acute graft-versus-host disease in adult and pediatric patients 12 years and older (1.3)

-----DOSAGE AND ADMINISTRATION ------

Doses should be individualized based on safety and efficacy. Starting doses per indication are noted below. Myelofibrosis (2.1)

- The starting dose of Jakafi is based on patient's baseline platelet count:
 - Greater than 200 X 10⁹/L: 20 mg given orally twice daily
 - $100 \times 10^9 / L$ to $200 \times 10^9 / L$: 15 mg given orally twice daily
 - 50×10^9 /L to less than 100×10^9 /L: 5 mg given orally twice daily
- Monitor complete blood counts every 2 to 4 weeks until doses are stabilized, and then as clinically indicated. Modify or interrupt dosing for thrombocytopenia.

Polycythemia Vera (2.2)

• The starting dose of Jakafi is 10 mg given orally twice daily.

Acute Graft-Versus-Host Disease (2.3)

• The starting dose of Jakafi is 5 mg given orally twice daily.

DOSAGE FORMS AND STRENGTHS
Tablets: 5 mg, 10 mg, 15 mg, 20 mg and 25 mg. (3)
Tablets. 5 mg, 10 mg, 15 mg, 20 mg and 25 mg. (5)
CONTRAINDICATIONS
None. (4)

- Thrombocytopenia, Anemia and Neutropenia: Manage by dose reduction, or interruption, or transfusion. (5.1)
- Risk of Infection: Assess patients for signs and symptoms of infection and initiate appropriate treatment promptly. Serious infections should have resolved before starting therapy with Jakafi. (5.2)
- Symptom Exacerbation Following Interruption or Discontinuation: Manage with supportive care and consider resuming treatment with Jakafi. (5.3)
- Risk of Non-Melanoma Skin Cancer: Perform periodic skin examinations. (5.4)
- Lipid Elevations: Assess lipid levels 8-12 weeks from start of therapy and treat as needed. (5.5)

----- ADVERSE REACTIONS -----

- In myelofibrosis and polycythemia vera, the most common hematologic adverse reactions (incidence > 20%) are thrombocytopenia and anemia. The most common nonhematologic adverse reactions (incidence $\geq 15\%$) are bruising, dizziness, headache, and diarrhea. (6.1 and 6.2)
- In acute graft-versus-host disease, the most common hematologic adverse reactions (incidence > 50%) are anemia, thrombocytopenia, and neutropenia. The most common nonhematologic adverse reactions (incidence > 50%) are infections and edema. (6.3)

To report SUSPECTED ADVERSE REACTIONS, contact Incyte Corporation at 1-855-463-3463 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch

----- DRUG INTERACTIONS ·····

• Strong CYP3A4 Inhibitors or Fluconazole: Reduce, interrupt, or discontinue Jakafi doses as recommended. Avoid use of Jakafi with fluconazole doses greater than 200 mg except in patients with acute graft-versus-host disease. (2.4, 7)

------USE IN SPECIFIC POPULATIONS ------

- Renal Impairment: Reduce Jakafi starting dose or avoid treatment as recommended. (2.5, 8.6)
- Hepatic Impairment: Reduce Jakafi starting dose or avoid treatment as recommended. (2.5, 8.7)
- Lactation: Advise not to breastfeed. (8.2)

See 17 for PATIENT COUNSELING INFORMATION and FDA-approved patient labeling.

Revised: 2/2020

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FULL PRESCRIBING INFORMATION

1. INDICATIONS AND USAGE

1.1 Myelofibrosis

Jakafi is indicated for treatment of intermediate or high-risk myelofibrosis (MF), including primary MF, post-polycythemia vera MF and post-essential thrombocythemia MF in adults.

1.2 Polycythemia Vera

Jakafi is indicated for treatment of polycythemia vera (PV) in adults who have had an inadequate response to or are intolerant of hydroxyurea.

1.3 Acute Graft-Versus-Host Disease

Jakafi is indicated for treatment of steroid-refractory acute graft-versus-host disease (GVHD) in adult and pediatric patients 12 years and older.

2. DOSAGE AND ADMINISTRATION

2.1 Myelofibrosis

The recommended starting dose of Jakafi is based on platelet count (Table 1). A complete blood count (CBC) and platelet count must be performed before initiating therapy, every 2 to 4 weeks until doses are stabilized, and then as clinically indicated [see Warnings and Precautions (5.1)]. Doses may be titrated based on safety and efficacy.

Table 1: Jakafi Starting Doses for Myelofibrosis

Platelet Count	Starting Dose
Greater than 200 X 10 ⁹ /L	20 mg orally twice daily
$100 \text{ X } 10^9\text{/L} \text{ to } 200 \text{ X } 10^9\text{/L}$	15 mg orally twice daily
50 X 10 ⁹ /L to less than 100 X 10 ⁹ /L	5 mg orally twice daily

Dose Modification Guidelines for Hematologic Toxicity for Patients with Myelofibrosis Starting Treatment with a Platelet Count of $100 \times 10^9/L$ or Greater

Treatment Interruption and Restarting Dosing

^{*} Sections or subsections omitted from the full prescribing information are not listed.

Interrupt treatment for platelet counts less than 50 X 10^9 /L or absolute neutrophil count (ANC) less than 0.5 X 10^9 /L.

After recovery of platelet counts above $50 \times 10^9/L$ and ANC above $0.75 \times 10^9/L$, dosing may be restarted. Table 2 illustrates the maximum allowable dose that may be used in restarting Jakafi after a previous interruption.

Table 2: Myelofibrosis: Maximum Restarting Doses for Jakafi after Safety Interruption for Thrombocytopenia for Patients Starting Treatment with a Platelet Count of 100 X $10^9/L$ or Greater

Current Platelet Count	Maximum Dose When Restarting Jakafi Treatment*
Greater than or equal to 125 X 10 ⁹ /L	20 mg twice daily
100 to less than 125 X 10 ⁹ /L	15 mg twice daily
75 to less than 100 X 10 ⁹ /L	10 mg twice daily for at least 2 weeks; if stable, may increase to 15 mg twice daily
50 to less than 75 X 10 ⁹ /L	5 mg twice daily for at least 2 weeks; if stable, may increase to 10 mg twice daily
Less than 50 X 10 ⁹ /L	Continue hold

^{*} Maximum doses are displayed. When restarting, begin with a dose at least 5 mg twice daily below the dose at interruption.

Following treatment interruption for ANC below 0.5×10^9 /L, after ANC recovers to 0.75×10^9 /L or greater, restart dosing at the higher of 5 mg once daily or 5 mg twice daily below the largest dose in the week prior to the treatment interruption.

Dose Reductions

Dose reductions should be considered if the platelet counts decrease as outlined in Table 3 with the goal of avoiding dose interruptions for thrombocytopenia.

Table 3: Myelofibrosis: Dosing Recommendations for Thrombocytopenia for Patients Starting Treatment with a Platelet Count of $100 \times 10^9/L$ or Greater

	Dose at Time of Platelet Decline				
Platelet Count	25 mg twice daily	20 mg twice daily	15 mg twice daily	10 mg twice daily	5 mg twice daily
	New Dose	New Dose	New Dose	New Dose	New Dose
100 to less than 125 X 10 ⁹ /L	20 mg twice daily	15 mg twice daily	No Change	No Change	No Change
75 to less than 100 X 10 ⁹ /L	10 mg twice daily	10 mg twice daily	10 mg twice daily	No Change	No Change
50 to less than	5 mg	5 mg	5 mg	5 mg	No

75 X 10 ⁹ /L	twice daily	twice daily	twice daily	twice daily	Change
Less than 50 X 10 ⁹ /L	Hold	Hold	Hold	Hold	Hold

Dose Modification Based on Insufficient Response for Patients with Myelofibrosis Starting Treatment with a Platelet Count of 100×10^9 /L or Greater

If the response is insufficient and platelet and neutrophil counts are adequate, doses may be increased in 5 mg twice daily increments to a maximum of 25 mg twice daily. Doses should not be increased during the first 4 weeks of therapy and not more frequently than every 2 weeks.

Consider dose increases in patients who meet all of the following conditions:

- a. Failure to achieve a reduction from pretreatment baseline in either palpable spleen length of 50% or a 35% reduction in spleen volume as measured by computed tomography (CT) or magnetic resonance imaging (MRI);
- b. Platelet count greater than 125 X 10⁹/L at 4 weeks and platelet count never below 100 X 10⁹/L;
- c. ANC Levels greater than 0.75×10^9 /L.

Based on limited clinical data, long-term maintenance at a 5 mg twice daily dose has not shown responses and continued use at this dose should be limited to patients in whom the benefits outweigh the potential risks. Discontinue Jakafi if there is no spleen size reduction or symptom improvement after 6 months of therapy.

Dose Modifications for Hematologic Toxicity for Patients with Myelofibrosis Starting Treatment with Platelet Counts of 50 \times 10 9 /L to Less Than 100 \times 10 9 /L

This section applies only to patients with platelet counts of 50×10^9 /L to less than 100×10^9 /L prior to any treatment with Jakafi. See dose modifications in Section 2.1 (*Dose Modification Guidelines for Hematological Toxicity for Patients with Myelofibrosis Starting Treatment with a Platelet Count of 100 \times 10^9/L or Greater) for hematological toxicity in patients whose platelet counts were 100 \times 10^9/L or more prior to starting treatment with Jakafi.*

Treatment Interruption and Restarting Dosing

Interrupt treatment for platelet counts less than 25 X $10^9/L$ or ANC less than 0.5 X $10^9/L$.

After recovery of platelet counts above 35×10^9 /L and ANC above 0.75×10^9 /L, dosing may be restarted. Restart dosing at the higher of 5 mg once daily or 5 mg twice daily below the largest dose in the week prior to the decrease in platelet count below 25×10^9 /L or ANC below 0.5×10^9 /L that led to dose interruption.

Dose Reductions

Reduce the dose of Jakafi for platelet counts less than 35 \times 10⁹/L as described in Table 4.

Table 4: Myelofibrosis: Dosing Modifications for Thrombocytopenia for Patients with Starting Platelet Count of 50 \times 10 9 /L to Less Than 100 \times 10 9 /L

Platelet Count	Dosing Recommendations
Less than 25 X 10 ⁹ /L	Interrupt dosing.
25 X 10 ⁹ /L to less than 35 X 10 ⁹ /L AND the platelet count decline is less than 20% during the prior four weeks	 Decrease dose by 5 mg once daily. For patients on 5 mg once daily, maintain dose at 5 mg once daily.
25×10^9 /L to less than 35×10^9 /L	Decrease dose by 5 mg twice daily.

AND the platelet count decline is 20% or greater during the prior four weeks

- For patients on 5 mg twice daily, decrease the dose to 5 mg once daily.
- For patients on 5 mg once daily, maintain dose at 5 mg once daily.

Dose Modifications Based on Insufficient Response for Patients with Myelofibrosis and Starting Platelet Count of 50 X 10⁹/L to Less Than 100 X 10⁹/L

Do not increase doses during the first 4 weeks of therapy, and do not increase the dose more frequently than every 2 weeks.

If the response is insufficient as defined in Section 2.1 (see Dose Modification Based on Insufficient Response with Myelofibrosis Starting Treatment with a platelet count of 100 X 10⁹/L or Greater), doses may be increased by increments of 5 mg daily to a maximum of 10 mg twice daily if:

- a. the platelet count has remained at least 40×10^9 /L, and
- b. the platelet count has not fallen by more than 20% in the prior 4 weeks, and
- c. the ANC is more than $1 \times 10^9/L$, and
- d. the dose has not been reduced or interrupted for an adverse event or hematological toxicity in the prior 4 weeks.

Continuation of treatment for more than 6 months should be limited to patients in whom the benefits outweigh the potential risks. Discontinue Jakafi if there is no spleen size reduction or symptom improvement after 6 months of therapy.

Dose Modification for Bleeding

Interrupt treatment for bleeding requiring intervention regardless of current platelet count. Once the bleeding event has resolved, consider resuming treatment at the prior dose if the underlying cause of bleeding has been controlled. If the bleeding event has resolved but the underlying cause persists, consider resuming treatment with Jakafi at a lower dose.

2.2 Polycythemia Vera

The recommended starting dose of Jakafi is 10 mg twice daily. Doses may be titrated based on safety and efficacy.

Dose Modification Guidelines for Patients with Polycythemia Vera

A complete blood count (CBC) and platelet count must be performed before initiating therapy, every 2 to 4 weeks until doses are stabilized, and then as clinically indicated [see Warnings and Precautions (5.1)].

Dose Reductions

Dose reductions should be considered for hemoglobin and platelet count decreases as described in Table 5.

Table 5: Polycythemia Vera: Dose Reductions

Hemoglobin and/or Platelet Count	Dosing Recommendations
Hemoglobin greater than or equal to $12~g/dL$ AND platelet count greater than or equal to $100~X~10^9/L$	No change required.
Hemoglobin 10 to less than 12 g/dL AND platelet count 75 to less than 100 X 10 ⁹ /L	• Dose reductions should be considered with the goal of avoiding dose interruptions for anemia and thrombocytopenia.

Hemoglobin 8 to less than 10 g/dL OR platelet count 50 to less than 75 X 10 ⁹ /L	 Reduce dose by 5 mg twice daily. For patients on 5 mg twice daily, decrease the dose to 5 mg once daily.
Hemoglobin less than 8 g/dL OR platelet count less than 50 X 10 ⁹ /L	Interrupt dosing.

Treatment Interruption and Restarting Dosing

Interrupt treatment for hemoglobin less than 8 g/dL, platelet counts less than 50 X 10^9 /L or ANC less than 1.0 X 10^9 /L.

After recovery of the hematologic parameter(s) to acceptable levels, dosing may be restarted.

Table 6 illustrates the dose that may be used in restarting Jakafi after a previous interruption.

Table 6: Polycythemia Vera: Restarting Doses for Jakafi after Safety Interruption for Hematologic Parameter(s)

Use the **most severe category** of a patient's hemoglobin, platelet count, or ANC abnormality to determine the corresponding maximum restarting dose.

Hemoglobin, Platelet Count, or ANC	Maximum Restarting Dose
Hemoglobin less than 8 g/dL OR	Continue hold
platelet count less than 50 X 10 ⁹ /L OR	
ANC less than 1 X 10 ⁹ /L	
	5 mg twice daily*or no more than
	5 mg twice daily less than the dose
ANC 1 to less than 1.5 X 10 ⁹ /L	which resulted in dose interruption
Hemoglobin 10 to less than 12 g/dL OR	10 mg twice daily*or no more than
	5 mg twice daily less than the dose
ANC 1.5 to less than 2 X 10 ⁹ /L	which resulted in dose interruption
Hemoglobin greater than or equal to 12 g/dL OR	15 mg twice daily*or no more than
platelet count greater than or equal to 100 X 10 ⁹ /L OR	5 mg twice daily less than the dose
ANC greater than or equal to 2 X 10 ⁹ /L	which resulted in dose interruption

^{*} Continue treatment for at least 2 weeks; if stable, may increase dose by 5 mg twice daily.

Patients who had required dose interruption while receiving a dose of 5 mg twice daily, may restart at a dose of 5 mg twice daily or 5 mg once daily, but not higher, once hemoglobin is greater than or equal to 10 g/dL, platelet count is greater than or equal to 75×10^9 /L, and ANC is greater than or equal to 1.5×10^9 /L.

Dose Management after Restarting Treatment

After restarting Jakafi following treatment interruption, doses may be titrated, but the maximum total daily dose should not exceed 5 mg less than the dose that resulted in the dose interruption. An exception to this is dose interruption following phlebotomy-associated anemia, in which case the maximal total daily dose allowed after restarting Jakafi would not be limited.

Dose Modifications Based on Insufficient Response for Patients with Polycythemia Vera

If the response is insufficient and platelet, hemoglobin, and neutrophil counts are adequate, doses may be increased in 5 mg twice daily increments to a maximum of 25 mg twice daily. Doses should not be increased during the first 4 weeks of therapy and not more frequently than every two weeks.

Consider dose increases in patients who meet all of the following conditions:

- 1. Inadequate efficacy as demonstrated by one or more of the following:
 - a. Continued need for phlebotomy
 - b. WBC greater than the upper limit of normal range
 - c. Platelet count greater than the upper limit of normal range
 - d. Palpable spleen that is reduced by less than 25% from Baseline
- 2. Platelet count greater than or equal to $140 \times 10^9/L$
- 3. Hemoglobin greater than or equal to 12 g/dL
- 4. ANC greater than or equal to $1.5 \times 10^9/L$

2.3 Acute Graft-Versus-Host Disease

The recommended starting dose of Jakafi is 5 mg given orally twice daily. Consider increasing the dose to 10 mg twice daily after at least 3 days of treatment if the ANC and platelet counts are not decreased by 50% or more relative to the first day of dosing with Jakafi.

Tapering of Jakafi may be considered after 6 months of treatment in patients with response who have discontinued therapeutic doses of corticosteroids. Taper Jakafi by one dose level approximately every 8 weeks (10 mg twice daily to 5 mg twice daily to 5 mg once daily). If acute GVHD signs or symptoms recur during or after the taper of Jakafi, consider retreatment.

Dose Modification Guidelines for Patients with Acute Graft-Versus-Host Disease

Evaluate blood parameters before and during treatment with Jakafi. Dose reductions should be considered for platelet counts, ANCs or bilirubin value as described in Table 7. Patients who are currently receiving Jakafi 10 mg twice daily may have their dose reduced to 5 mg twice daily; patients receiving 5 mg twice daily may have their dose reduced to 5 mg once daily. Patients who are unable to tolerate Jakafi at a dose of 5 mg once daily should have treatment interrupted until their clinical and/or laboratory parameters recover.

Table 7: Dose Modifications for Patients with Acute GVHD

Laboratory Parameter	Dosing Recommendations
Clinically significant	Reduce dose by 1 dose level.
thrombocytopenia after supportive	When platelets recover to previous values, dosing may
measures	return to prior dose level.
ANC less than 1 X 10 ⁹ /L	Hold Jakafi for up to 14 days; resume at 1 dose level
considered related to Jakafi	lower upon recovery.
	$3.0-5.0 \times \text{ULN}$: Continue Jakafi at 1 dose level lower
	until recovery.
	$>5.0-10.0 \times ULN$: Hold Jakafi for up to 14 days until
•	bilirubin ≤ 1.5 × ULN; resume at current dose upon
GVHD	recovery
	Total bilirubin > 10.0 × ULN: Hold Jakafi for up to
	14 days until bilirubin ≤ 1.5 × ULN; resume at 1 dose
	level lower upon recovery.
Total Bilirubin elevation, liver	>3.0 × ULN: Continue Jakafi at 1 dose level lower
GVHD	until recovery.

2.4 Dose Modifications for Concomitant Use with Strong CYP3A4 Inhibitors or Fluconazole

Modify the Jakafi dosage when coadministered with strong CYP3A4 inhibitors and fluconazole doses of less than or equal to 200 mg [see Drug Interactions (7)], according to Table 8.

Additional dose modifications should be made with frequent monitoring of safety and efficacy.

Avoid the use of fluconazole doses of greater than 200 mg daily with Jakafi except in patients with acute GVHD.

Table 8: Dose Modifications for Concomitant Use with Strong CYP3A4 Inhibitors or Fluconazole

For patients coadministered strong CYP3A4 inhibitors or fluconazole doses of less than or equal to 200 mg	Recommended Dose Modification		
Starting dose for patients with MF with a platelet count:			
• Greater than or equal to 100 X 10 ⁹ /L	10 mg twice daily		
• 50 X 10 ⁹ /L to less than 100 X 10 ⁹ /L	5 mg once daily		
Starting dose for patients with PV:	5 mg twice daily		
If on stable dose for patients with MF or PV:			
Greater than or equal to 10 mg twice daily	Decrease dose by 50% (round up to the closest available tablet strength)		
• 5 mg twice daily	5 mg once daily		
• 5 mg once daily	Avoid strong CYP3A4 inhibitor or fluconazole treatment or interrupt Jakafi treatment for the duration of strong CYP3A4 inhibitor or fluconazole use		
For patients with acute GVHD:			
Ketoconazole	5 mg once daily		
Other CYP3A4 inhibitors*	No dose adjustment		

^{*} With coadministration of itraconazole, monitor blood counts more frequently for toxicity and adjust the dose of Jakafi if necessary.

2.5 Dose Modifications for Renal or Hepatic Impairment

Renal Impairment

Patients with Moderate or Severe Renal Impairment

Modify the Jakafi dosage for patients with moderate or severe renal impairment according to Table 9.

Patients with End Stage Renal Disease on Dialysis

Modify the Jakafi dosage for patients with end stage renal disease (ESRD) on dialysis according to Table 9. Make additional dose modifications with frequent monitoring of safety and efficacy. Avoid use of Jakafi in patients with ESRD (CLcr less than 15 mL/min) not requiring dialysis [see Use in Specific Populations (8.6)].

Table 9: Dose Modifications for Renal Impairment

Renal Impairment Status	Platelet Count	Recommended Starting Dosage
Patients with MF		
	Greater than 150 X 10 ⁹ /L	No dose modification needed
	100 + 150 V 109л	10 ma turing daily

Moderate (CLcr 30 to 59 mL/min)	100 to 130 V 10-/F	TO HIS IMICE UALLY
or	50 to less than 100 X	5 mg daily
Severe (CLcr 15 to 29 mL/min)	$10^{9}/L$	5 filg dairy
Severe (CECI 13 to 23 IIIL/IIIIII)	Less than 50 X 10 ⁹ /L	Avoid use [see Use in Specific
	Less than 50 X 10°/L	Populations (8.6)]
ESRD (CLcr less than 15 mL/min)	100 to 200 X 10 ⁹ /L	15 mg once after dialysis session
on	Greater than 200 X 10 ⁹ /L	20 mg once often dialysis session
dialysis	Greater than 200 X 10°/L	20 mg once after dialysis session
Patients with PV		
Moderate (CLcr 30 to 59 mL/min)		
or	Any	5 mg twice daily
Severe (CLcr 15 to 29 mL/min)		
ESRD (CLcr less than 15 mL/min)		
on	Any	10 mg once after dialysis session
dialysis		
Patients with acute GVHD		
Moderate (CLcr 30 to 59 mL/min)		
or	Any	5 mg once daily
Severe (CLcr 15 to 29 mL/min)		
ESRD (CLcr less than 15 mL/min)		
on	Any	5 mg once after dialysis session
dialysis		

ESRD = end stage renal disease, and CLcr = creatinine clearance

Hepatic Impairment

Modify the Jakafi dosage for patients with hepatic impairment according to Table 10.

Table 10: Dose Modifications for Hepatic Impairment

Hepatic Impairment Status	Platelet Count	Recommended Starting Dosage
	Greater than 150 X 10 ⁹ /L	No dose modification needed
Patients with MF	100×10^9 /L to 150×10^9 /L	10 mg twice daily
Mild, Moderate, or Severe (Child-	50 to less than 100 X 10 ⁹ /L	5 mg daily
Pugh Class A, B, C)	Less than 50 X 10 ⁹ /L	Avoid use [see Use in Specific Populations (8.7)]
Patients with PV		
Mild, Moderate, or Severe (Child-Pugh Class A, B, C)	Any	5 mg twice daily
Patients with acute GVHD		
Mild, Moderate, or Severe based on NCI criteria	Any	No dose modification needed
		Monitor blood counts more
Stage 3 or 4 liver GVHD	Any	frequently for toxicity and
		consider 5 mg once daily

2.6 Method of Administration

Jakafi is dosed orally and can be administered with or without food.

If a dose is missed, the patient should not take an additional dose, but should take the next usual prescribed dose.

When discontinuing Jakafi therapy for reasons other than thrombocytopenia, gradual tapering of the dose of Jakafi may be considered, for example by 5 mg twice daily each week.

For patients unable to ingest tablets, Jakafi can be administered through a nasogastric tube (8 French or greater) as follows:

- Suspend one tablet in approximately 40 mL of water with stirring for approximately 10 minutes.
- Within 6 hours after the tablet has dispersed, the suspension can be administered through a nasogastric tube using an appropriate syringe.

The tube should be rinsed with approximately 75 mL of water. The effect of tube feeding preparations on Jakafi exposure during administration through a nasogastric tube has not been evaluated.

3. DOSAGE FORMS AND STRENGTHS

5 mg tablets - round and white with "INCY" on one side and "5" on the other.

10 mg tablets - round and white with "INCY" on one side and "10" on the other.

15 mg tablets - oval and white with "INCY" on one side and "15" on the other.

20 mg tablets - capsule-shaped and white with "INCY" on one side and "20" on the other.

25 mg tablets - oval and white with "INCY" on one side and "25" on the other.

4. CONTRAINDICATIONS

None.

5. WARNINGS AND PRECAUTIONS

5.1 Thrombocytopenia, Anemia and Neutropenia

Treatment with Jakafi can cause thrombocytopenia, anemia and neutropenia [see Dosage and Administration (2.1)].

Manage thrombocytopenia by reducing the dose or temporarily interrupting Jakafi. Platelet transfusions may be necessary [see Dosage and Administration (2), and Adverse Reactions (6.1)].

Patients developing anemia may require blood transfusions and/or dose modifications of Jakafi.

Severe neutropenia (ANC less than 0.5×10^9 /L) was generally reversible by withholding Jakafi until recovery [see Adverse Reactions (6.1)].

Perform a pre-treatment complete blood count (CBC) and monitor CBCs every 2 to 4 weeks until doses are stabilized, and then as clinically indicated [see Dosage and Administration (2), and Adverse Reactions (6.1)].

5.2 Risk of Infection

Serious bacterial, mycobacterial, fungal and viral infections have occurred. Delay starting therapy with Jakafi until active serious infections have resolved. Observe patients receiving Jakafi for signs and symptoms of infection and manage promptly. Use active surveillance and prophylactic antibiotics according to clinical guidelines.

Tuberculosis

Tuberculosis infection has been reported in patients receiving Jakafi. Observe patients receiving Jakafi for signs and symptoms of active tuberculosis and manage promptly.

Prior to initiating Jakafi, patients should be evaluated for tuberculosis risk factors, and those at higher

risk should be tested for latent infection. Risk factors include, but are not limited to, prior residence in or travel to countries with a high prevalence of tuberculosis, close contact with a person with active tuberculosis, and a history of active or latent tuberculosis where an adequate course of treatment cannot be confirmed.

For patients with evidence of active or latent tuberculosis, consult a physician with expertise in the treatment of tuberculosis before starting Jakafi. The decision to continue Jakafi during treatment of active tuberculosis should be based on the overall risk-benefit determination.

Progressive Multifocal Leukoencephalopathy

Progressive multifocal leukoencephalopathy (PML) has occurred with Jakafi treatment. If PML is suspected, stop Jakafi and evaluate.

Herpes Zoster

Advise patients about early signs and symptoms of herpes zoster and to seek treatment as early as possible if suspected [see Adverse Reactions (6.1)].

Hepatitis B

Hepatitis B viral load (HBV-DNA titer) increases, with or without associated elevations in alanine aminotransferase and aspartate aminotransferase, have been reported in patients with chronic HBV infections taking Jakafi. The effect of Jakafi on viral replication in patients with chronic HBV infection is unknown. Patients with chronic HBV infection should be treated and monitored according to clinical guidelines.

5.3 Symptom Exacerbation Following Interruption or Discontinuation of Treatment with Jakafi

Following discontinuation of Jakafi, symptoms from myeloproliferative neoplasms may return to pretreatment levels over a period of approximately one week. Some patients with MF have experienced one or more of the following adverse events after discontinuing Jakafi: fever, respiratory distress, hypotension, DIC, or multi-organ failure. If one or more of these occur after discontinuation of, or while tapering the dose of Jakafi, evaluate for and treat any intercurrent illness and consider restarting or increasing the dose of Jakafi. Instruct patients not to interrupt or discontinue Jakafi therapy without consulting their physician. When discontinuing or interrupting therapy with Jakafi for reasons other than thrombocytopenia or neutropenia [see Dosage and Administration (2.6)], consider tapering the dose of Jakafi gradually rather than discontinuing abruptly.

5.4 Non-Melanoma Skin Cancer

Non-melanoma skin cancers including basal cell, squamous cell, and Merkel cell carcinoma have occurred in patients treated with Jakafi. Perform periodic skin examinations.

5.5 Lipid Elevations

Treatment with Jakafi has been associated with increases in lipid parameters including total cholesterol, low-density lipoprotein (LDL) cholesterol, and triglycerides. The effect of these lipid parameter elevations on cardiovascular morbidity and mortality has not been determined in patients treated with Jakafi. Assess lipid parameters approximately 8-12 weeks following initiation of Jakafi therapy. Monitor and treat according to clinical guidelines for the management of hyperlipidemia.

6. ADVERSE REACTIONS

The following clinically significant adverse reactions are discussed in greater detail in other sections of the labeling:

- Thrombocytopenia, Anemia and Neutropenia [see Warnings and Precautions (5.1)]
- Risk of Infection [see Warnings and Precautions (5.2)]
- Symptom Exacerbation Following Interruption or Discontinuation of Treatment with Jakafi [see

Warnings and Precautions (5.3)

• Non-Melanoma Skin Cancer [see Warnings and Precautions (5.4)]

6.1 Clinical Trials Experience in Myelofibrosis

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect the rates observed in practice.

The safety of Jakafi was assessed in 617 patients in six clinical studies with a median duration of follow-up of 10.9 months, including 301 patients with MF in two Phase 3 studies.

In these two Phase 3 studies, patients had a median duration of exposure to Jakafi of 9.5 months (range 0.5 to 17 months), with 89% of patients treated for more than 6 months and 25% treated for more than 12 months. One hundred and eleven (111) patients started treatment at 15 mg twice daily and 190 patients started at 20 mg twice daily. In patients starting treatment with 15 mg twice daily (pretreatment platelet counts of 100 to 200 X 10^9 /L) and 20 mg twice daily (pretreatment platelet counts greater than 200 X 10^9 /L), 65% and 25% of patients, respectively, required a dose reduction below the starting dose within the first 8 weeks of therapy.

In a double-blind, randomized, placebo-controlled study of Jakafi, among the 155 patients treated with Jakafi, the most frequent adverse reactions were thrombocytopenia and anemia [see Table 12]. Thrombocytopenia, anemia and neutropenia are dose-related effects. The three most frequent nonhematologic adverse reactions were bruising, dizziness and headache [see Table 11].

Discontinuation for adverse events, regardless of causality, was observed in 11% of patients treated with Jakafi and 11% of patients treated with placebo.

Table 11 presents the most common nonhematologic adverse reactions occurring in patients who received Jakafi in the double-blind, placebo-controlled study during randomized treatment.

Table 11: Myelofibrosis: Nonhematologic Adverse Reactions Occurring in Patients on Jakafi in the Double-blind, Placebo-controlled Study During Randomized Treatment

	Jakafi (N=155)			Placebo (N=151)		
Adverse Reactions	All Grades* (%)	Grade 3 (%)	Grade 4 (%)	All Grades (%)	Grade 3 (%)	Grade 4 (%)
Bruising [†]	23	<1	0	15	0	0
Dizziness [‡]	18	<1	0	7	0	0
Headache	15	0	0	5	0	0
Urinary Tract Infections§	9	0	0	5	<1	<1
Weight Gain [¶]	7	<1	0	1	<1	0
Flatulence	5	0	0	<1	0	0
Herpes Zoster [#]	2	0	0	<1	0	0

^{*} National Cancer Institute Common Terminology Criteria for Adverse Events (CTCAE), version 3.0

[†] includes contusion, ecchymosis, hematoma, injection site hematoma, periorbital hematoma, vessel puncture site hematoma, increased tendency to bruise, petechiae, purpura

[‡] includes dizziness, postural dizziness, vertigo, balance disorder, Meniere's Disease, labyrinthitis

[§] includes urinary tract infection, cystitis, urosepsis, urinary tract infection bacterial, kidney infection, pyuria, bacteria urine, bacteria urine identified, nitrite urine present

[¶] includes weight increased, abnormal weight gain

[#] includes herpes zoster and post-herpetic neuralgia

Description of Selected Adverse Reactions

Anemia

In the two Phase 3 clinical studies, median time to onset of first CTCAE Grade 2 or higher anemia was approximately 6 weeks. One patient (<1%) discontinued treatment because of anemia. In patients receiving Jakafi, mean decreases in hemoglobin reached a nadir of approximately 1.5 to 2.0 g/dL below baseline after 8 to 12 weeks of therapy and then gradually recovered to reach a new steady state that was approximately 1.0 g/dL below baseline. This pattern was observed in patients regardless of whether they had received transfusions during therapy.

In the randomized, placebo-controlled study, 60% of patients treated with Jakafi and 38% of patients receiving placebo received red blood cell transfusions during randomized treatment. Among transfused patients, the median number of units transfused per month was 1.2 in patients treated with Jakafi and 1.7 in placebo treated patients.

Thrombocytopenia

In the two Phase 3 clinical studies, in patients who developed Grade 3 or 4 thrombocytopenia, the median time to onset was approximately 8 weeks. Thrombocytopenia was generally reversible with dose reduction or dose interruption. The median time to recovery of platelet counts above 50 X 10^9 /L was 14 days. Platelet transfusions were administered to 5% of patients receiving Jakafi and to 4% of patients receiving control regimens. Discontinuation of treatment because of thrombocytopenia occurred in <1% of patients receiving Jakafi and <1% of patients receiving control regimens. Patients with a platelet count of 100 X 10^9 /L to 200 X 10^9 /L before starting Jakafi had a higher frequency of Grade 3 or 4 thrombocytopenia compared to patients with a platelet count greater than 200 X 10^9 /L (17% versus 7%).

Neutropenia

In the two Phase 3 clinical studies, 1% of patients reduced or stopped Jakafi because of neutropenia.

Table 12 provides the frequency and severity of clinical hematology abnormalities reported for patients receiving treatment with Jakafi or placebo in the placebo-controlled study.

Table 12: Myelofibrosis: Worst Hematology Laboratory Abnormalities in the
Placebo-Controlled Study*

	Jakafi (N=155)				Placebo (N=151)	
	All Grades † Grade 3 Grade 4			All Grades	Grade 3	Grade 4
Laboratory Parameter	(%)	(%)	(%)	(%)	(%)	(%)
Thrombocytopenia	70	9	4	31	1	0
Anemia	96	34	11	87	16	3
Neutropenia	19	5	2	4	<1	1

^{*} Presented values are worst Grade values regardless of baseline

Additional Data from the Placebo-Controlled Study

- 25% of patients treated with Jakafi and 7% of patients treated with placebo developed newly occurring or worsening Grade 1 abnormalities in alanine transaminase (ALT). The incidence of greater than or equal to Grade 2 elevations was 2% for Jakafi with 1% Grade 3 and no Grade 4 ALT elevations.
- 17% of patients treated with Jakafi and 6% of patients treated with placebo developed newly
 occurring or worsening Grade 1 abnormalities in aspartate transaminase (AST). The incidence of

[†] National Cancer Institute Common Terminology Criteria for Adverse Events, version 3.0

Grade 2 AST elevations was <1% for Jakafi with no Grade 3 or 4 AST elevations.

• 17% of patients treated with Jakafi and <1% of patients treated with placebo developed newly occurring or worsening Grade 1 elevations in cholesterol. The incidence of Grade 2 cholesterol elevations was <1% for Jakafi with no Grade 3 or 4 cholesterol elevations.

6.2 Clinical Trial Experience in Polycythemia Vera

In a randomized, open-label, active-controlled study, 110 patients with PV resistant to or intolerant of hydroxyurea received Jakafi and 111 patients received best available therapy [see Clinical Studies (14.2)]. The most frequent adverse reaction was anemia. Discontinuation for adverse events, regardless of causality, was observed in 4% of patients treated with Jakafi. Table 13 presents the most frequent nonhematologic adverse reactions occurring up to Week 32.

Table 13: Polycythemia Vera: Nonhematologic Adverse Reactions Occurring in ≥ 5% of Patients on Jakafi in the Open-Label, Active-controlled Study up to Week 32 of Randomized Treatment

	Jak (N=1		Best Available Therapy (N=111)		
	All Grades*	Grade 3-4	All Grades	Grade 3-4	
Adverse Reactions	(%)	(%)	(%)	(%)	
Diarrhea	15	0	7	<1	
Dizziness [†]	15	0	13	0	
Dyspnea [‡]	13	3	4	0	
Muscle Spasms	12	<1	5	0	
Constipation	8	0	3	0	
Herpes Zoster [§]	6	<1	0	0	
Nausea	6	0	4	0	
Weight Gain [¶]	6	0	<1	0	
Urinary Tract Infections#	6	0	3	0	
Hypertension	5	<1	3	<1	

^{*} National Cancer Institute Common Terminology Criteria for Adverse Events (CTCAE), version 3.0

Clinically relevant laboratory abnormalities are shown in Table 14.

Table 14: Polycythemia Vera: Selected Laboratory Abnormalities in the Open-Label, Active-controlled Study up to Week 32 of Randomized Treatment*

	Jakafi (N=110)			Best A	vailable ' (N=111)	10
	All			All		
	Grades [†]	Grade 3	Grade 4	Grades	Grade 3	Grade 4
Laboratory Parameter	(%)	(%)	(%)	(%)	(%)	(%)
Hematology						
Anemia	72	<1	<1	58	0	0

[†] includes dizziness and vertigo

[‡] includes dyspnea and dyspnea exertional

[§] includes herpes zoster and post-herpetic neuralgia

 $[\]P$ includes weight increased and abnormal weight gain

[#] includes urinary tract infection and cystitis

Thrombocytopenia	27	5	<1	24	3	<1
Neutropenia	3	0	<1	10	<1	0
Chemistry						
Hypercholesterolemia	35	0	0	8	0	0
Elevated ALT	25	<1	0	16	0	0
Elevated AST	23	0	0	23	<1	0
Hypertriglyceridemia	15	0	0	13	0	0

^{*} Presented values are worst Grade values regardless of baseline

6.3 Clinical Trial Experience in Acute Graft-Versus-Host Disease

In a single-arm, open-label study, 71 adults (ages 18-73 years) were treated with Jakafi for acute GVHD failing treatment with steroids with or without other immunosuppressive drugs [see *Clinical Studies*(14.3)]. The median duration of treatment with Jakafi was 46 days (range, 40382 days).

There were no fatal adverse reactions to Jakafi. An adverse reaction resulting in treatment discontinuation occurred in 31% of patients. The most common adverse reaction leading to treatment discontinuation was infection (10%). Table 15 shows the adverse reactions other than laboratory abnormalities.

Table 15: Acute Graft-Versus-Host Disease:
Nonhematologic Adverse Reactions Occurring in ≥
15% of Patients in the Open-Label, Single-Cohort
Study

	Jakafi (N	N=71)
Adverse Reactions*	All Grades [†] (%)	Grade 3-4 (%)
Infections	55	41
Edema	51	13
Hemorrhage	49	20
Fatigue	37	14
Bacterial infections	32	28
Dyspnea	32	7
Viral infections	31	14
Thrombosis	25	11
Diarrhea	24	7
Rash	23	3
Headache	21	4
Hypertension	20	13
Dizziness	16	0

^{*} Selected laboratory abnormalities are listed in Table 16 below

Selected laboratory abnormalities during treatment with Jakafi are shown in Table 16.

[†] National Cancer Institute Common Terminology Criteria for Adverse Events, version 3.0

[†] National Cancer Institute Common Terminology Criteria for Adverse Events (CTCAE), version 4.03

	Jakafi (N=71)					
	Worst grade during treatmen					
Laboratory Parameter	All Grades* (%)	Grade 3-4 (%)				
Hematology						
Anemia	75	45				
Thrombocytopenia	75	61				
Neutropenia	58	40				
Chemistry						
Elevated ALT	48	8				
Elevated AST	48	6				
Hypertriglyceridemia	11	1				

^{*} National Cancer Institute Common Terminology Criteria for Adverse Events, version 4.03

7. DRUG INTERACTIONS

Fluconazole

Concomitant administration of Jakafi with fluconazole doses greater than 200 mg daily may increase ruxolitinib exposure due to inhibition of both the CYP3A4 and CYP2C9 metabolic pathways [see Clinical Pharmacology (12.3)]. Increased exposure may increase the risk of exposure-related adverse reactions. Avoid the concomitant use of Jakafi with fluconazole doses of greater than 200 mg daily except in patients with acute GVHD [see Dosage and Administration (2.4)].

Strong CYP3A4 inhibitors

Concomitant administration of Jakafi with strong CYP3A4 inhibitors increases ruxolitinib exposure [see Clinical Pharmacology (12.3)]. Increased exposure may increase the risk of exposure-related adverse reactions. Consider dose reduction when administering Jakafi with strong CYP3A4 inhibitors [see Dosage and Administration (2.4)]. In patients with acute GVHD, reduce Jakafi dose as recommended only when coadministered with ketoconazole, and monitor blood counts more frequently for toxicity and adjust the dose if necessary when coadministered with itraconazole [see Dosage and Administration (2.4)].

Strong CYP3A4 inducers

Concomitant administration of Jakafi with strong CYP3A4 inducers may decrease ruxolitinib exposure [see Clinical Pharmacology (12.3)]. No dose adjustment is recommended; however, monitor patients frequently and adjust the Jakafi dose based on safety and efficacy [see Clinical Pharmacology (12.3)].

8. USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summary

When pregnant rats and rabbits were administered ruxolitinib during the period of organogenesis adverse developmental outcomes occurred at doses associated with maternal toxicity (*see Data*). There are no studies with the use of Jakafi in pregnant women to inform drug-associated risks.

The background risk of major birth defects and miscarriage for the indicated populations is unknown. Adverse outcomes in pregnancy occur regardless of the health of the mother or the use of medications. The background risk in the U.S. general population of major birth defects is 2% to 4% and miscarriage is 15% to 20% of clinically recognized pregnancies.

Data

Animal Data

Ruxolitinib was administered orally to pregnant rats or rabbits during the period of organogenesis, at doses of 15, 30 or 60 mg/kg/day in rats and 10, 30 or 60 mg/kg/day in rabbits. There were no treatment-related malformations. Adverse developmental outcomes, such as decreases of approximately 9% in fetal weights were noted in rats at the highest and maternally toxic dose of 60 mg/kg/day. This dose results in an exposure (AUC) that is approximately 2 times the clinical exposure at the maximum recommended dose of 25 mg twice daily. In rabbits, lower fetal weights of approximately 8% and increased late resorptions were noted at the highest and maternally toxic dose of 60 mg/kg/day. This dose is approximately 7% the clinical exposure at the maximum recommended dose.

In a pre- and post-natal development study in rats, pregnant animals were dosed with ruxolitinib from implantation through lactation at doses up to 30 mg/kg/day. There were no drug-related adverse findings in pups for fertility indices or for maternal or embryofetal survival, growth and development parameters at the highest dose evaluated (34% the clinical exposure at the maximum recommended dose of 25 mg twice daily).

8.2 Lactation

Risk Summary

No data are available regarding the presence of ruxolitinib in human milk, the effects on the breast fed child, or the effects on milk production. Ruxolitinib and/or its metabolites were present in the milk of lactating rats (*see Data*). Because many drugs are present in human milk and because of the potential for thrombocytopenia and anemia shown for Jakafi in human studies, discontinue breastfeeding during treatment with Jakafi and for two weeks after the final dose.

Data

Animal Data

Lactating rats were administered a single dose of [¹⁴C]-labeled ruxolitinib (30 mg/kg) on postnatal Day 10, after which plasma and milk samples were collected for up to 24 hours. The AUC for total radioactivity in milk was approximately 13-fold the maternal plasma AUC. Additional analysis showed the presence of ruxolitinib and several of its metabolites in milk, all at levels higher than those in maternal plasma.

8.4 Pediatric Use

The safety and effectiveness of Jakafi for treatment of myelofibrosis or polycythemia vera in pediatric patients have not been established.

The safety and effectiveness of Jakafi for treatment of steroid-refractory acute graft-versus-host disease (GVHD) have been established for treatment of children 12 years and older. Use of Jakafi in pediatric patients with steroid-refractory acute GVHD is supported by evidence from an adequate and well-controlled trial of Jakafi in adults [see Clinical Studies (14.3)] and additional pharmacokinetic and safety data in pediatric patients.

Jakafi was evaluated in a single-arm, dose-escalation study (NCT01164163) in 27 pediatric patients with relapsed or refractory solid tumors (Cohort A) and 20 with leukemias or myeloproliferative neoplasms (Cohort B). The patients had a median age of 14 years (range, 2 to 21 years) and included 18 children (age 2 to <12 years), and 14 adolescents (age 12 to <17 years). The dose levels tested were 15, 21, 29, 39, or 50 mg/m² twice daily in 28-day cycles with up to 6 patients per dose group.

Overall, 38 (81%) patients were treated with no more than a single cycle of Jakafi, while 3, 1, 2, and 3 patients received 2, 3, 4, and 5 or more cycles, respectively. A protocol-defined maximal tolerated dose was not observed, but since few patients were treated for multiple cycles, tolerability with continued use was not assessed adequately to establish a recommended Phase 2 dose higher than the recommended

dose for adults. The safety profile in children was similar to that seen in adults.

Juvenile Animal Toxicity Data

Administration of ruxolitinib to juvenile rats resulted in effects on growth and bone measures. When administered starting at postnatal day 7 (the equivalent of a human newborn) at doses of 1.5 to 75 mg/kg/day, evidence of fractures occurred at doses \geq 30 mg/kg/day, and effects on body weight and other bone measures [e.g., bone mineral content, peripheral quantitative computed tomography, and x-ray analysis] occurred at doses \geq 5 mg/kg/day. When administered starting at postnatal day 21 (the equivalent of a human 2-3 years of age) at doses of 5 to 60 mg/kg/day, effects on body weight and bone occurred at doses \geq 15 mg/kg/day, which were considered adverse at 60 mg/kg/day. Males were more severely affected than females in all age groups, and effects were generally more severe when administration was initiated earlier in the postnatal period. These findings were observed at exposures that are at least 27% the clinical exposure at the maximum recommended dose of 25 mg twice daily.

8.5 Geriatric Use

Of the total number of patients with MF in clinical studies with Jakafi, 52% were 65 years and older, while 15% were 75 years and older. No overall differences in safety or effectiveness of Jakafi were observed between these patients and younger patients.

Clinical studies of Jakafi in patients with acute GVHD did not include sufficient numbers of subjects age 65 and over to determine whether they respond differently from younger subjects.

8.6 Renal Impairment

Total exposure of ruxolitinib and its active metabolites increased with moderate (CLcr 30 mL/min to 59 mL/min) and severe (CLcr 15 mL/min to 29 mL/min) renal impairment, and ESRD on dialysis [see Clinical Pharmacology (12.3)]. Reduce Jakafi dose as recommended [see Dosage and Administration (2.5)].

8.7 Hepatic Impairment

Exposure of ruxolitinib increased with mild (Child-Pugh A), moderate (Child-Pugh B) and severe (Child-Pugh C) hepatic impairment [see Clinical Pharmacology (12.3)].

Reduce Jakafi dose as recommended in patients with MF or PV and any hepatic impairment [see Dosage and Administration (2.5)].

Monitor blood counts more frequently for toxicity and consider 5 mg once daily for patients with Stage 3 or 4 liver GVHD [see Dosage and Administration (2.5) and Clinical Pharmacology (12.3)].

10. OVERDOSAGE

There is no known antidote for overdoses with Jakafi. Single doses up to 200 mg have been given with acceptable acute tolerability. Higher than recommended repeat doses are associated with increased myelosuppression including leukopenia, anemia and thrombocytopenia. Appropriate supportive treatment should be given.

Hemodialysis is not expected to enhance the elimination of Jakafi.

11. DESCRIPTION

Ruxolitinib phosphate is a kinase inhibitor with the chemical name (*R*)-3-(4-(7*H*-pyrrolo[2,3-*d*]pyrimidin-4-yl)-1*H*-pyrazol-1-yl)-3-cyclopentylpropanenitrile phosphate and a molecular weight of 404.36. Ruxolitinib phosphate has the following structural formula:

Ruxolitinib phosphate is a white to off-white to light pink powder and is soluble in aqueous buffers across a pH range of 1 to 8.

Jakafi (ruxolitinib) Tablets are for oral administration. Each tablet contains ruxolitinib phosphate equivalent to 5 mg, 10 mg, 15 mg, 20 mg and 25 mg of ruxolitinib free base together with microcrystalline cellulose, lactose monohydrate, magnesium stearate, colloidal silicon dioxide, sodium starch glycolate, povidone and hydroxypropyl cellulose.

12. CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Ruxolitinib, a kinase inhibitor, inhibits Janus Associated Kinases (JAKs) JAK1 and JAK2 which mediate the signaling of a number of cytokines and growth factors that are important for hematopoiesis and immune function. JAK signaling involves recruitment of STATs (signal transducers and activators of transcription) to cytokine receptors, activation and subsequent localization of STATs to the nucleus leading to modulation of gene expression.

MF and PV are myeloproliferative neoplasms (MPN) known to be associated with dysregulated JAK1 and JAK2 signaling. In a mouse model of JAK2V617F-positive MPN, oral administration of ruxolitinib prevented splenomegaly, preferentially decreased JAK2V617F mutant cells in the spleen and decreased circulating inflammatory cytokines (e.g., $TNF-\alpha$, IL-6).

JAK-STAT signaling pathways play a role in regulating the development, proliferation, and activation of several immune cell types important for GVHD pathogenesis. In a mouse model of acute GVHD, oral administration of ruxolitinib was associated with decreased expression of inflammatory cytokines in colon homogenates and reduced immune-cell infiltration in the colon.

12.2 Pharmacodynamics

Jakafi inhibits cytokine induced STAT3 phosphorylation in whole blood from patients with MF and PV. Jakafi administration resulted in maximal inhibition of STAT3 phosphorylation 2 hours after dosing which returned to near baseline by 10 hours in patients with MF and PV.

Cardiac Electrophysiology

At a dose of 1.25 to 10 times the highest recommended starting dosage, Jakafi does not prolong the QT interval to any clinically relevant extent.

12.3 Pharmacokinetics

Mean ruxolitinib maximal plasma concentration (C_{max}) and AUC increased proportionally over a single dose range of 5 mg to 200 mg. Mean ruxolitinib C_{max} ranged from 205 nM to 7100 nM and AUC ranged from 862 nM*hr to 30700 nM*hr over a single dose range of 5 mg to 200 mg.

Absorption

Ruxolitinib achieves C_{max} within 1 hour to 2 hours post-dose. Oral absorption of ruxolitinib is estimated to be at least 95%.

Food Effect

No clinically relevant changes in the pharmacokinetics of ruxolitinib were observed upon administration of Jakafi with a high-fat, high-calorie meal (approximately 800 to 1000 calories of which 50% were derived from fat).

Distribution

The mean volume of distribution at steady-state is 72 L (coefficient of variation [CV] 29%) in patients with MF and 75 L (23%) in patients with PV.

Binding to plasma proteins is approximately 97%, mostly to albumin.

Elimination

The mean elimination half-life of ruxolitinib is approximately 3 hours and the mean half-life of ruxolitinib + metabolites is approximately 5.8 hours.

Ruxolitinib clearance (% coefficient of variation, CV) was 17.7 L/h in women and 22.1 L/h in men with MF (39%).

Ruxolitinib clearance (%CV) was 12.7 L/h (42%) in patients with PV.

Ruxolitinib clearance (%CV) was 11.9 L/h (43%) in patients with acute GVHD.

Metabolism

Ruxolitinib is metabolized by CYP3A4 and to a lesser extent by CYP2C9.

Excretion

Following a single oral dose of radiolabeled ruxolitinib, elimination was predominately through metabolism with 74% of radioactivity excreted in urine and 22% excretion via feces. Unchanged drug accounted for less than 1% of the excreted total radioactivity.

Specific Populations

No clinically relevant differences in ruxolitinib pharmacokinetics were observed with regard to age, race, sex, or weight. No clinically relevant effect in ruxolitinib pharmacokinetics were observed with regards to any hepatic impairment (total bilirubin >ULN and any aspartate transferase) in patients with acute GVHD.

Patients with Renal Impairment

Following oral administration of a single dose of Jakafi 25 mg, the total AUC of ruxolitinib and its active metabolites increased by 1.3-,1.5-, and 1.9-fold in subjects with mild, moderate, and severe renal impairment, respectively, compared to that in subjects with normal renal function (CLcr \geq 90 mL/min). Also, the total AUC of ruxolitinib and its active metabolites increased by 1.6-fold in subjects with ESRD after dialysis) compared to that in subjects with normal renal function (CLcr \geq 90 mL/min). The pharmacokinetics of ruxolitinib was similar in subjects with various degrees of renal impairment and in those with normal renal function. The change in the pharmacodynamic marker, pSTAT3 inhibition, was consistent with the corresponding increase in metabolite exposure with renal impairment. Ruxolitinib is not removed by dialysis; however, the removal of some active metabolites by dialysis cannot be ruled out.

Patients with Hepatic Impairment

Following oral administration of a single dose of Jakafi 25 mg, the AUC of ruxolitinib increased in subjects with mild (Child-Pugh A) by 1.9-fold, moderate (Child-Pugh B) by 1.3-fold, and severe (Child-Pugh C) hepatic impairment by 1.7-fold compared to that in subjects with normal hepatic function.

The change in the pharmacodynamic marker, pSTAT3 inhibition, was consistent with the corresponding increase in ruxolitinib exposure except in the severe hepatic impairment cohort where the pharmacodynamic activity was more prolonged in some subjects than expected based on plasma

concentrations of ruxolitinib.

Drug Interactions

Fluconazole

Simulations suggest that fluconazole (a dual CYP3A4 and CYP2C9 inhibitor) increases steady state ruxolitinib AUC by approximately 100% to 300% following concomitant administration of 10 mg of Jakafi twice daily with 100 mg to 400 mg of fluconazole once daily [see Dosage and Administration (2.4) and Drug Interactions (7)].

Strong CYP3A4 inhibitors

Ketoconazole (a strong CYP3A4 inhibitor) increased ruxolitinib C_{max} by 33% and AUC by 91%. Ketoconazole also prolonged ruxolitinib half-life from 3.7 hours to 6 hours [see Dosage and Administration (2.4) and Drug Interactions (7)].

Moderate CYP3A4 inhibitors

Erythromycin (a moderate CYP3A4 inhibitor) increased ruxolitinib C_{max} by 8% and AUC by 27% [see *Drug Interactions (7)*].

Strong CYP3A4 inducers

Rifampin (a strong CYP3A4 inducer) decreased ruxolitinib C_{max} by 32% and AUC by 61%. The relative exposure to ruxolitinib's active metabolites increased approximately 100% [see *Drug Interactions (7)*].

In vitro studies

Ruxolitinib and its M18 metabolite did not inhibit CYP1A2, CYP2B6, CYP2C8, CYP2C9, CYP2C19, CYP2D6 or CYP3A4. Ruxolitinib did not induce CYP1A2, CYP2B6 or CYP3A4 at clinically relevant concentrations.

Ruxolitinib and its M18 metabolite did not inhibit the P-gp, BCRP, OATP1B1, OATP1B3, OCT1, OCT2, OAT1 or OAT3 transport systems at clinically relevant concentrations. Ruxolitinib is not a substrate for the P-gp transporter.

13. NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Ruxolitinib was not carcinogenic in the 6-month Tg.rasH2 transgenic mouse model or in a 2-year carcinogenicity study in the rat.

Ruxolitinib was not mutagenic in a bacterial mutagenicity assay (Ames test) or clastogenic in *in vitro* chromosomal aberration assay (cultured human peripheral blood lymphocytes) or *in vivo* in a rat bone marrow micronucleus assay.

In a fertility study, ruxolitinib was administered to male rats prior to and throughout mating and to female rats prior to mating and up to the implantation day (gestation day 7). Ruxolitinib had no effect on fertility or reproductive function in male or female rats at doses of 10, 30 or 60 mg/kg/day. However, in female rats doses of greater than or equal to 30 mg/kg/day resulted in increased post-implantation loss. The exposure (AUC) at the dose of 30 mg/kg/day is approximately 34% the clinical exposure at the maximum recommended dose of 25 mg twice daily.

14. CLINICAL STUDIES

14.1 Myelofibrosis

Two randomized Phase 3 studies (Studies 1 and 2) were conducted in patients with MF (either primary

MF, post-polycythemia vera MF or post-essential thrombocythemia-MF). In both studies, patients had palpable splenomegaly at least 5 cm below the costal margin and risk category of intermediate 2 (2 prognostic factors) or high risk (3 or more prognostic factors) based on the International Working Group Consensus Criteria (IWG).

The starting dose of Jakafi was based on platelet count. Patients with a platelet count between 100 and 200 X 10^9 /L were started on Jakafi 15 mg twice daily and patients with a platelet count greater than 200 X 10^9 /L were started on Jakafi 20 mg twice daily. Doses were then individualized based upon tolerability and efficacy with maximum doses of 20 mg twice daily for patients with platelet counts between 100 to less than or equal to 125 X 10^9 /L, of 10 mg twice daily for patients with platelet counts between 75 to less than or equal to 100×10^9 /L, and of 5 mg twice daily for patients with platelet counts between 50 to less than or equal to 75×10^9 /L.

Study 1

Study 1 (NCT00952289) was a double-blind, randomized, placebo-controlled study in 309 patients who were refractory to or were not candidates for available therapy. The median age was 68 years (range 40 to 91 years) with 61% of patients older than 65 years and 54% were male. Fifty percent (50%) of patients had primary MF, 31% had post-polycythemia vera MF and 18% had post-essential thrombocythemia MF. Twenty-one percent (21%) of patients had red blood cell transfusions within 8 weeks of enrollment in the study. The median hemoglobin count was 10.5 g/dL and the median platelet count was 251 X 10⁹/L. Patients had a median palpable spleen length of 16 cm below the costal margin, with 81% having a spleen length 10 cm or greater below the costal margin. Patients had a median spleen volume as measured by magnetic resonance imaging (MRI) or computed tomography (CT) of 2595 cm³ (range 478 cm³ to 8881 cm³). (The upper limit of normal is approximately 300 cm³).

Patients were dosed with Jakafi or matching placebo. The primary efficacy endpoint was the proportion of patients achieving greater than or equal to a 35% reduction from baseline in spleen volume at Week 24 as measured by MRI or CT.

Secondary endpoints included duration of a 35% or greater reduction in spleen volume and proportion of patients with a 50% or greater reduction in Total Symptom Score from baseline to Week 24 as measured by the modified Myelofibrosis Symptom Assessment Form (MFSAF) v2.0 diary.

Study 2

Study 2 (NCT00934544) was an open-label, randomized study in 219 patients. Patients were randomized 2:1 to Jakafi versus best available therapy. Best available therapy was selected by the investigator on a patient-by-patient basis. In the best available therapy arm, the medications received by more than 10% of patients were hydroxyurea (47%) and glucocorticoids (16%). The median age was 66 years (range 35 to 85 years) with 52% of patients older than 65 years and 57% were male. Fifty-three percent (53%) of patients had primary MF, 31% had post-polycythemia vera MF and 16% had post-essential thrombocythemia MF. Twenty-one percent (21%) of patients had red blood cell transfusions within 8 weeks of enrollment in the study. The median hemoglobin count was 10.4 g/dL and the median platelet count was 236 X 10⁹/L. Patients had a median palpable spleen length of 15 cm below the costal margin, with 70% having a spleen length 10 cm or greater below the costal margin. Patients had a median spleen volume as measured by MRI or CT of 2381 cm³ (range 451 cm³ to 7765 cm³).

The primary efficacy endpoint was the proportion of patients achieving 35% or greater reduction from baseline in spleen volume at Week 48 as measured by MRI or CT.

A secondary endpoint in Study 2 was the proportion of patients achieving a 35% or greater reduction of spleen volume as measured by MRI or CT from baseline to Week 24.

Study 1 and 2 Efficacy Results

Efficacy analyses of the primary endpoint in Studies 1 and 2 are presented in Table 17 below. A significantly larger proportion of patients in the Jakafi group achieved a 35% or greater reduction in spleen volume from baseline in both studies compared to placebo in Study 1 and best available therapy

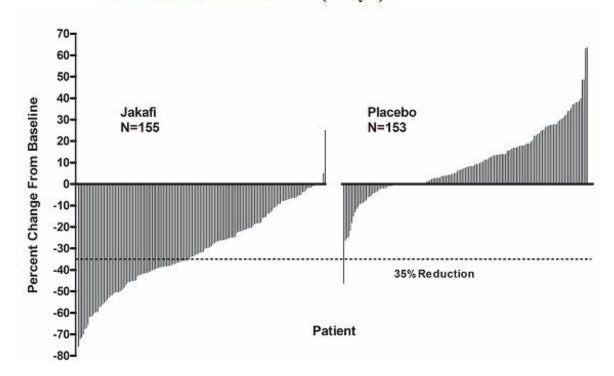
in Study 2. A similar proportion of patients in the Jakafi group achieved a 50% or greater reduction in palpable spleen length.

Table 17: Percent of Patients with Myelofibrosis Achieving 35% or Greater Reduction from Baseline in Spleen Volume at Week 24 in Study 1 and at Week 48 in Study 2 (Intent to Treat)

	Study 1 Jakafi Placebo (N=155) (N=154)		Study 2	
			Jakafi (N=146)	Best Available Therapy (N=73)
Time Points	Week 24		Week 48	
Number (%) of Patients with Spleen Volume Reduction by 35% or More	65 (42)	1 (<1)	41 (29)	0
P-value	< 0.0001		< 0.0001	

Figure 1 shows the percent change from baseline in spleen volume for each patient at Week 24 (Jakafi N=139, placebo N=106) or the last evaluation prior to Week 24 for patients who did not complete 24 weeks of randomized treatment (Jakafi N=16, placebo N=47). One (1) patient (placebo) with a missing baseline spleen volume is not included.

Figure 1: Percent Change from Baseline in Spleen Volume at Week 24 or Last Observation for Each Patient (Study 1)



In Study 1, MF symptoms were a secondary endpoint and were measured using the modified Myelofibrosis Symptom Assessment Form (MFSAF) v2.0 diary. The modified MFSAF is a daily diary capturing the core symptoms of MF (abdominal discomfort, pain under left ribs, night sweats, itching, bone/muscle pain and early satiety). Symptom scores ranged from 0 to 10 with 0 representing symptoms "absent" and 10 representing "worst imaginable" symptoms. These scores were added to create the

daily total score, which has a maximum of 60.

Table 18 presents assessments of Total Symptom Score from baseline to Week 24 in Study 1 including the proportion of patients with at least a 50% reduction (ie, improvement in symptoms). At baseline, the mean Total Symptom Score was 18.0 in the Jakafi group and 16.5 in the placebo group. A higher proportion of patients in the Jakafi group had a 50% or greater reduction in Total Symptom Score than in the placebo group, with a median time to response of less than 4 weeks.

Table 18: Improvement in Total Symptom Score in Patients with Myelofibrosis

	Jakafi (N=148)	Placebo (N=152)
Number (%) of Patients with 50% or Greater Reduction in Total Symptom Score by Week 24	68 (46)	8 (5)
P-value	< 0.00	001

Figure 2 shows the percent change from baseline in Total Symptom Score for each patient at Week 24 (Jakafi N=129, placebo N=103) or the last evaluation on randomized therapy prior to Week 24 for patients who did not complete 24 weeks of randomized treatment (Jakafi N=16, placebo N=42). Results are excluded for 5 patients with a baseline Total Symptom Score of zero, 8 patients with missing baseline and 6 patients with insufficient post-baseline data.

Figure 2: Percent Change from Baseline in Total Symptom Score at Week 24 or Last Observation for Each Patient (Study 1)

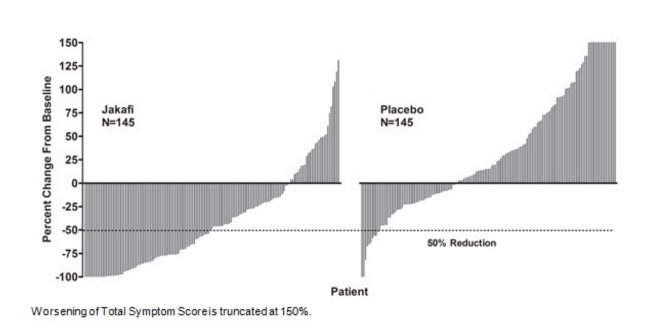
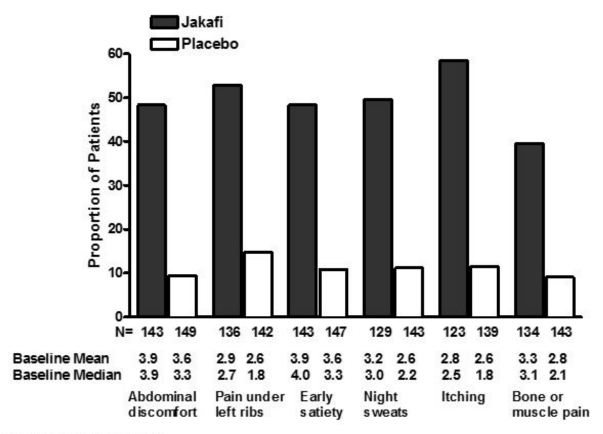


Figure 3 displays the proportion of patients with at least a 50% improvement in each of the individual symptoms that comprise the Total Symptom Score indicating that all 6 of the symptoms contributed to the higher Total Symptom Score response rate in the group treated with Jakafi.

Figure 3: Proportion of Patients with Myelofibrosis Achieving 50% or Greater Reduction in Individual Symptom Scores at Week 24



Individual score range = 0 to 10

An exploratory analysis of patients receiving Jakafi also showed improvement in fatigue-related symptoms (i.e., tiredness, exhaustion, mental tiredness, and lack of energy) and associated impacts on daily activities (i.e., activity limitations related to work, self-care, and exercise) as measured by the PROMIS[®] Fatigue 7-item short form total score at Week 24. Patients who achieved a reduction of 4.5 points or more from baseline to Week 24 in the PROMIS[®] Fatigue total score were considered to have achieved a fatigue response. Fatigue response was reported in 35% of patients in the Jakafi group versus 14% of the patients in the placebo group.

Overall survival was a secondary endpoint in both Study 1 and Study 2. Patients in the control groups were eligible for crossover in both studies, and the median times to crossover were 9 months in Study 1 and 17 months in Study 2.

Figure 4 and Figure 5 show Kaplan-Meier curves of overall survival at prospectively planned analyses after all patients remaining on study had completed 144 weeks on study.

Figure 4: Overall Survival - Kaplan-Meier Curves by Treatment Group in Study 1

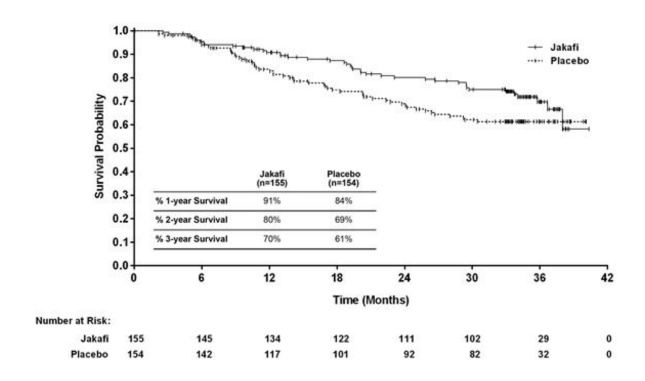
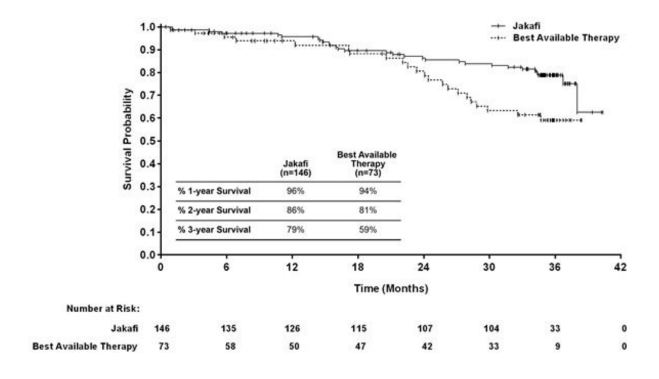


Figure 5: Overall Survival - Kaplan-Meier Curves by Treatment Group in Study 2



14.2 Polycythemia Vera

Study 3 (NCT01243944) was a randomized, open-label, active-controlled Phase 3 study conducted in

222 patients with PV. Patients had been diagnosed with PV for at least 24 weeks, had an inadequate response to or were intolerant of hydroxyurea, required phlebotomy and exhibited splenomegaly. All patients were required to demonstrate hematocrit control between 40-45% prior to randomization. The age ranged from 33 to 90 years with 30% of patients over 65 years of age and 66% were male. Patients had a median spleen volume as measured by MRI or CT of 1272 cm³ (range 254 cm³ to 5147 cm³) and median palpable spleen length below the costal margin was 7 cm.

Patients were randomized to Jakafi or best available therapy. The starting dose of Jakafi was 10 mg twice daily. Doses were then individualized based upon tolerability and efficacy with a maximum dose of 25 mg twice daily. At Week 32, 98 patients were still on Jakafi with 8% receiving greater than 20 mg twice daily, 15% receiving 20 mg twice daily, 33% receiving 15 mg twice daily, 34% receiving 10 mg twice daily, and 10% receiving less than 10 mg twice daily. Best available therapy (BAT) was selected by the investigator on a patient-by-patient basis and included hydroxyurea (60%), interferon/pegylated interferon (12%), anagrelide (7%), pipobroman (2%), lenalidomide/thalidomide (5%), and observation (15%).

The primary endpoint was the proportion of subjects achieving a response at Week 32, with response defined as having achieved both hematocrit control (the absence of phlebotomy eligibility beginning at the Week 8 visit and continuing through Week 32) and spleen volume reduction (a greater than or equal to 35% reduction from baseline in spleen volume at Week 32). Phlebotomy eligibility was defined as a confirmed hematocrit greater than 45% that is at least 3 percentage points higher than the hematocrit obtained at baseline or a confirmed hematocrit greater than 48%, whichever was lower. Secondary endpoints included the proportion of all randomized subjects who achieved the primary endpoint and who maintained their response 48 weeks after randomization, and the proportion of subjects achieving complete hematological remission at Week 32 with complete hematological remission defined as achieving hematocrit control, platelet count less than or equal to 400 X 10⁹/L, and white blood cell count less than or equal to 10 X 10⁹/L.

Results of the primary and secondary endpoints are presented in Table 19. A significantly larger proportion of patients on the Jakafi arm achieved a response for the primary endpoint compared to best available therapy at Week 32 and maintained their response 48 weeks after randomization. A significantly larger proportion of patients on the Jakafi arm compared to best available therapy also achieved complete hematological remission at Week 32.

Table 19: Percent of Patients with Polycythemia Vera Achieving the Primary and Key Secondary Endpoints in Study 3 (Intent to Treat)

	Jakafi (N=110)	Best Available Therapy (N=112)
Number (%) of Patients Achieving a Primary Response at Week 32	25 (23%)	1 (<1%)
95% CI of the response rate (%)	(15%, 32%)	(0%, 5%)
P-value	< 0.0	0001
Number (%) of Patients Achieving a Durable Primary Response at Week 48	22 (20%)	1 (<1%)
95% CI of the response rate (%)	(13%, 29%)	(0%, 5%)
P-value	< 0.0	0001
Number (%) of Patients Achieving Complete	GC (DAO/)	0 (00/1

Hematological Remission at Week 32	20 (24%)	y (০% <i>)</i>
95% CI of the response rate (%)	(16%, 33%)	(4%, 15%)
P-value	0.0	016

Primary Response defined as having achieved both the absence of phlebotomy eligibility beginning at the Week 8 visit and continuing through Week 32 and a greater than or equal to 35% reduction from baseline in spleen volume at Week 32.

Additional analyses for Study 3 to assess durability of response were conducted at Week 80 only in the Jakafi arm. On this arm, 91 (83%) patients were still on treatment at the time of the Week 80 data cut-off. Of the 25 patients who achieved a primary response at Week 32, 19 (76% of the responders) maintained their response through Week 80, and of the 26 patients who achieved complete hematological remission at Week 32, 15 (58% of the responders) maintained their response through Week 80.

In an assessment of the individual components that make up the primary endpoint, there were 66 (60%) patients with hematocrit control on the Jakafi arm vs. 21 (19%) patients on best available therapy at Week 32; 51 (77% of hematocrit responders) patients on the Jakafi arm maintained hematocrit control through Week 80. There were 44 (40%) patients with spleen volume reduction from baseline greater than or equal to 35% on the Jakafi arm vs. 1 (<1%) patient on best available therapy at Week 32; 43 (98% of spleen volume reduction responders) patients on the Jakafi arm maintained spleen volume reduction through Week 80.

14.3 Acute Graft-Versus-Host Disease

Study 4 (NCT02953678) was an open-label, single-arm, multicenter study of Jakafi for treatment of patients with steroid-refractory acute GVHD Grades 2 to 4 (Mount Sinai Acute GVHD International Consortium (MAGIC) criteria) occurring after allogeneic hematopoietic stem cell transplantation. Jakafi was administered at 5 mg twice daily, and the dose could be increased to 10 mg twice daily after 3 days in the absence of toxicity.

There were 49 patients with acute GVHD refractory to steroids alone. These patients had a median age of 57 years (range, 18-72 years), 47% were male, 92% were Caucasian, and 14% were Hispanic. At baseline, acute GVHD was Grade 2 in 27%, Grade 3 in 55%, and Grade 4 in 18%; 84% had visceral GVHD; the median MAGIC biomarker score was 0.47 (range, 0.10\(\text{0} \) 0.92); and the median ST2 level was 334 mcg/L (range, 55-1286 mcg/L). The median duration of prior corticosteroid exposure at baseline was 15 days (range: 3 - 106 days).

The efficacy of Jakafi was based on Day-28 overall response rate (ORR) (complete response, very good partial response or partial response by Center for International Blood and Marrow Transplant Research (CIBMTR) criteria) and the duration of response. The ORR results are presented in Table 20; Day-28 ORR was 100% for Grade 2 GVHD, 40.7% for Grade 3 GVHD, and 44.4% for Grade 4 GVHD. The median duration of response, calculated from Day-28 response to progression, new salvage therapy for acute GVHD or death from any cause (with progression being defined as worsening by one stage in any organ without improvement in other organs in comparison to prior response assessment) was 16 days (95% CI 9, 83). Also for the Day-28 responders, the median time from Day-28 response to either death or need for new therapy for acute GVHD (additional salvage therapy or increase in steroids) was 173 days (95% CI 66, NE).

Table 20: Day-28 Overall Response Rate for Patients with Steroid-Refractory Acute GVHD in Study 4

	Alone (n=49)
Overall Response (%) (95% CI)	28 (57.1%) (42.2, 71.2)
Complete Response	15 (30.6%)
Very Good Partial Response	2 (4.1%)
Partial Response	11 (22.4%)

16. HOW SUPPLIED/STORAGE AND HANDLING

Jakafi (ruxolitinib) Tablets are available as follows:

Jakafi Trade Presentations

NDC Number	Strength	Description	Tablets per Bottle
50881-005-60	5 mg	Round tablet with "INCY" on one side and "5" on the other	60
50881-010-60	10 mg	Round tablet with "INCY" on one side and "10" on the other	60
50881-015-60	15 mg	Oval tablet with "INCY" on one side and "15" on the other	60
50881-020-60	20 mg	Capsule shaped tablet with "INCY" on one side and "20" on the other	60
50881-025-60	25 mg	Oval tablet with "INCY" on one side and "25" on the othe	60

Store at room temperature 20°C to 25°C (68°F to 77°F); excursions permitted between 15°C and 30°C (59°F and 86°F) [see USP Controlled Room Temperature].

17. PATIENT COUNSELING INFORMATION

See FDA-approved patient labeling (Patient Information).

Discuss the following with patients prior to and during treatment with Jakafi:

Thrombocytopenia, Anemia and Neutropenia

Inform patients that Jakafi is associated with thrombocytopenia, anemia and neutropenia, and of the need to monitor complete blood counts before and during treatment. Advise patients to observe for and report bleeding.

Infections

Inform patients of the signs and symptoms of infection and to report any such signs and symptoms promptly.

Inform patients regarding the early signs and symptoms of herpes zoster and of progressive multifocal leukoencephalopathy, and advise patients to seek advice of a clinician if such symptoms are observed.

Symptom Exacerbation Following Interruption or Discontinuation of Treatment with Jakafi

Inform patients that after discontinuation of treatment, signs and symptoms from myeloproliferative neoplasms are expected to return. Instruct patients not to interrupt or discontinue Jakafi therapy without consulting their physician.

Non-Melanoma Skin Cancer

Inform patients that Jakafi may increase their risk of certain non-melanoma skin cancers. Advise patients to inform their healthcare provider if they have ever had any type of skin cancer or if they observe any new or changing skin lesions.

Lipid Elevations

Inform patients that Jakafi may increase blood cholesterol, and of the need to monitor blood cholesterol levels.

Drug-drug Interactions

Advise patients to inform their healthcare providers of all medications they are taking, including over-the-counter medications, herbal products and dietary supplements.

Dialysis

Inform patients on dialysis that their dose should not be taken before dialysis but only following dialysis.

Lactation

Inform women not to breastfeed during treatment with Jakafi and for two weeks after the final dose

Compliance

Advise patients to continue taking Jakafi every day for as long as their physician tells them and that this is a long-term treatment. Patients should not change dose or stop taking Jakafi without first consulting their physician. Patients should be aware that after discontinuation of treatment, signs and symptoms from myeloproliferative neoplasms are expected to return.

Manufactured for:

Incyte Corporation 1801 Augustine Cut-off Wilmington, DE 19803

Jakafi is a registered trademark of Incyte. All rights reserved.

U.S. Patent Nos. 7598257; 8415362; 8722693; 8822481; 8829013; 9079912; 9814722; 10016429 © 2011-2020 Incyte Corporation. All rights reserved.

Patient Information JAKAFI® (JAK-ah-fye)

(ruxolitinib) tablets

What is Jakafi?

Jakafi is a prescription medicine used to treat:

- adults with certain types of myelofibrosis (MF).
- adults with polycythemia vera (PV) who have already taken a medicine called hydroxyurea and it did not work well enough or they could not tolerate it
- adults and children 12 years of age and older with acute graft versus host disease (GVHD) who have taken corticosteroids and they did not work well enough.

It is not known if Jakafi is safe or effective in children for treatment of myelofibrosis or polycythemia vera.

Before taking Jakafi, tell your healthcare provider about of your medical conditions, including if you:

- have an infection
- have or had tuberculosis (TB), or have been in close contact with someone who has TB
- have or had hepatitis B

- have or have had liver problems
- have or have had kidney problems or are on dialysis. If you are on dialysis, Jakafi should be taken after your dialysis
- have high level of fat in your blood (high blood cholesterol or triglycerides)
- have had skin cancer in the past
- are pregnant or plan to become pregnant. It is not known if Jakafi will harm your unborn baby.
- are breastfeeding or plan to breastfeed. It is not known if Jakafi passes into your breast milk. Do not breastfeed during treatment with Jakafi and for 2 weeks after the final dose.

Tell your healthcare provider about all the medicines you take, including prescription and over-the-counter medicines, vitamins and herbal supplements. Taking Jakafi with certain other medicines may affect how Jakafi works. Know the medicines you take. Keep a list of them to show your healthcare provider and pharmacist when you get a new medicine.

How should I take Jakafi?

- Take Jakafi exactly as your healthcare provider tells you.
- Do not change your dose or stop taking Jakafi without first talking to your healthcare provider.
- You can take Jakafi with or without food.
- Jakafi may also be given through certain nasogastric tubes.
 - Tell your healthcare provider if you cannot take Jakafi by mouth. Your healthcare provider will decide if you can take Jakafi through a nasogastric tube.
 - Ask your healthcare provider to give you specific instruction on how to properly take Jakafi through a nasogastric tube.
- If you miss a dose of Jakafi, take your next dose at your regular time. Do not take 2 doses at the same time.
- If you take too much Jakafi call your healthcare provider or go to the nearest hospital emergency room right away.
- You will have regular blood tests during your treatment with Jakafi. Your healthcare provider may change your dose of Jakafi or stop your treatment based on the results of your blood tests.

What are the possible side effects of Jakafi?

Jakafi can cause serious side effects including:

Low blood cell counts. Jakafi may cause low platelet counts (thrombocytopenia), low red blood cell counts (anemia), and low white blood cell counts (neutropenia). If you develop bleeding, stop Jakafi and call your healthcare provider. Your healthcare provider will do a blood test to check your blood cell counts before you start Jakafi and regularly during your treatment with Jakafi. Tell your healthcare provider right away if you develop or have worsening of any of these symptoms:

- unusual bleeding
- bruising
- tiredness

- shortness of breath
- fever

Infection. You may be at risk for developing a serious infection during treatment with Jakafi. Tell your healthcare provider if you develop any of the following symptoms of infection:

- chills
- aches
- fever
- nausea

- vomiting
- weakness
- painful skin rash or blisters

Skin cancers. Some people who take Jakafi have developed certain types of non-melanoma skin cancers. Tell your healthcare provider if you develop any new or changing skin lesions during

treatment with Jakafi.

Choles terol increases. You may have changes in your blood cholesterol levels. Your healthcare provider will do blood tests to check your cholesterol levels during treatment with Jakafi.

The most common side effects of Jakafi in adults with certain types of MF and PV include:

- low platelet counts (thrombocytopenia)
- low red blood cell counts (anemia)
- bruising

- dizziness
- headache
- diarrhea

The most common side effects of Jakafi in people with acute graft versus host disease (GVHD) include:

- low red blood cell counts (anemia)
- low platelet counts (thrombocytopenia)
- low white blood cell counts (neutropenia)

- infections
- fluid retention

These are not all the possible side effects of Jakafi.

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

You may also report side effects to Incyte Corporation at 1-855-463-3463.

How should I store Jakafi?

• Store Jakafi at room temperature 68°F to 77°F (20°C to 25°C).

Keep Jakafi and all medicines out of the reach of children.

General information about the safe and effective use of Jakafi.

Medicines are sometimes prescribed for purposes other than those listed in Patient Information. Do not use Jakafi for a condition for which it is not prescribed. Do not give Jakafi to other people, even if they have the same symptoms you have. It may harm them. You can ask your pharmacist or healthcare provider for information that is written for healthcare professionals.

What are the ingredients in Jakafi?

Active ingredient: ruxolitinib phosphate

Inactive ingredients: microcrystalline cellulose, lactose monohydrate, magnesium stearate, colloidal silicon dioxide, sodium starch glycolate, povidone and hydroxypropyl cellulose

Manufactured for: Incyte Corporation, 1801 Augustine Cut-off, Wilmington, DE 19803

Jakafi is a registered trademark of Incyte. All rights reserved.

U.S. Patent Nos. 7598257; 8415362; 8722693; 8822481; 8829013; 9079912; 9814722; 10016429

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For more information call 1-855-463-3463 or go to www.jakafi.com.

This Patient Information has been approved by the U.S. Food and Drug Administration. Revised: January 2020

5 mg Tablet Bottle Label

Rx only

NDC 50881-005-60

Jakafi® (Ruxolitinib) Tablets

5 mg

60 tablets

Each tablet contains ruxolitinib phosphate equivalent to 5 mg ruxolitinib free base.



10 mg Tablet Bottle Label

Rx only

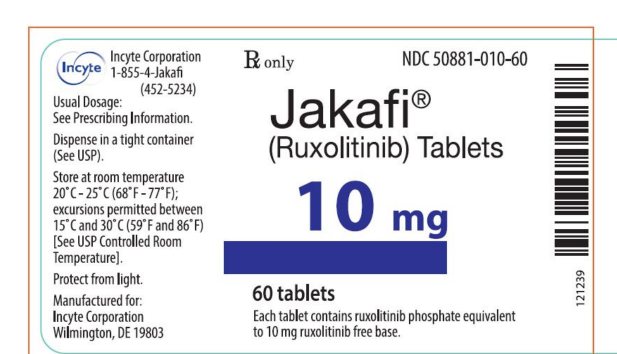
NDC 50881-010-60

Jakafi® (Ruxolitinib) Tablets

10 mg

60 tablets

Each tablet contains ruxolitinib phosphate equivalent to 10 mg ruxolitinib free base.



15 mg Tablet Bottle Label

Rx only

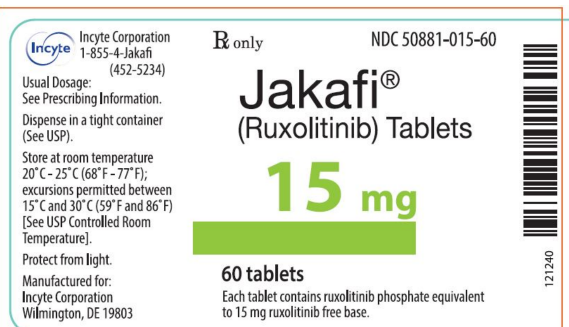
NDC 50881-015-60

Jakafi® (Ruxolitinib) Tablets

15 mg

60 tablets

Each tablet contains ruxolitinib phosphate equivalent to 15 mg ruxolitinib free base.



20 mg Tablet Bottle Label

Rx only

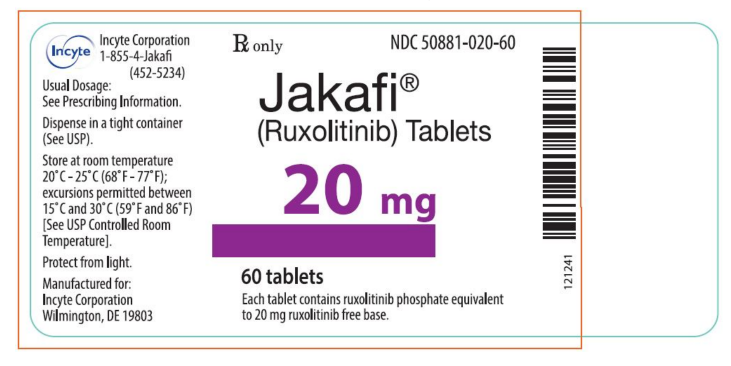
NDC 50881-020-60

Jakafi® (Ruxolitinib) Tablets

20 mg

60 tablets

Each tablet contains ruxolitinib phosphate equivalent to 20 mg ruxolitinib free base.



25 mg Tablet Bottle Label

Rx only

NDC 50881-025-60

Jakafi® (Ruxolitinib) Tablets

25 mg

60 tablets

Each tablet contains ruxolitinib phosphate equivalent to 25 mg ruxolitinib free base.



Usual Dosage: See Prescribing Information.

Dispense in a tight container (See USP).

Store at room temperature 20°C - 25°C (68°F - 77°F); excursions permitted between 15°C and 30°C (59°F and 86°F) [See USP Controlled Room Temperature].

Protect from light.

Manufactured for: Incyte Corporation Wilmington, DE 19803 Ronly

NDC 50881-025-60

Jakafi® (Ruxolitinib) Tablets

25 mg



60 tablets

Each tablet contains ruxolitinib phosphate equivalent to 25 mg ruxolitinib free base.

21242

JAKAFI

ruxolitinib tablet

Product Information

Product TypeHUMAN PRESCRIPTION DRUGItem Code (Source)NDC:50881-005

Route of Administration ORAL

Active Ingredient/Active Moiety

Ingredient Name
Basis of Strength
RUXOLITINIB (UNII: 82S8X8XX8H) (RUXOLITINIB - UNII:82S8X8XX8H)
RUXOLITINIB 5.0 mg

Product Characteristics

Color	WHITE (White to off-white)	Score	no score
Shape	ROUND	Size	7mm
Flavor		Imprint Code	INCY;5
Contains			

Packaging

# Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:50881-005-	60 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	11/16/2011	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA202192	11/16/2011	

JAKAFI

ruxolitinib tablet

Product	Information
I I VUULL	IIIIVI IIIA UVII

Route of Administration ORAL

Active Ingredient/Active Moiety

0	3		
	Ingredient Name	Basis of Strength	Strength
RUXOLITINIB (UN	NII: 82S8X8XX8H) (RUXOLITINIB - UNII:82S8X8XX8H)	RUXOLITINIB	10.0 mg

Inactive Ingredients	
Ingredient Name	Strength
CELLULO SE, MICRO CRYSTALLINE (UNII: OP1R32D61U)	
LACTO SE MONO HYDRATE (UNII: EWQ57Q8I5X)	
SILICON DIO XIDE (UNII: ETJ7Z6 XBU4)	
HYDROXYPROPYL CELLULOSE (70000 WAMW) (UNII: 6607AQV0RT)	
PO VIDO NE (UNII: FZ989 GH94E)	
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	

Product Characteristics				
Color	WHITE (White to off-white)	Score	no score	
Shape	ROUND	Size	9 mm	
Flavor		Imprint Code	INCY;10	
Contains				

Packaging#Item CodePackage DescriptionMarketing Start DateMarketing End Date1NDC:50881-010- 0128 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product11/16/20112NDC:50881-010- 6060 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product11/16/2011

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA202192	11/16/2011	

JAKAFI

ruxolitinib tablet

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:50881-015
Route of Administration	ORAL		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
RUXOLITINIB (UNII: 82S8X8XX8H) (RUXOLITINIB - UNII:82S8X8XX8H)	RUXOLITINIB	15.0 mg

Inactive Ingredients	
Ingredient Name	Strength
CELLULO SE, MICRO CRYSTALLINE (UNII: OP1R32D61U)	
LACTO SE MO NO HYDRATE (UNII: EWQ57Q8I5X)	
SILICON DIO XIDE (UNII: ETJ7Z6 XBU4)	
HYDROXYPROPYL CELLULOSE (70000 WAMW) (UNII: 6607AQV0RT)	
PO VIDO NE (UNII: FZ989 GH9 4E)	
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	

Product Characteristics				
Color	WHITE (White to off-white)	Score	no score	
Shape	OVAL	Size	15mm	
Flavor		Imprint Code	INCY;15	
Contains				

l	Pacl	kaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1 ND 60	C:50881-015-	60 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	11/16/2011	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA202192	11/16/2011	

JAKAFI

ruxolitinib tablet

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:50881-020
Route of Administration	ORAL		

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
RUXOLITINIB (UNII: 82S8X8XX8H) (RUXOLITINIB - UNII:82S8X8XX8H)	RUXOLITINIB	20.0 mg		

Inactive Ingredients	
Ingredient Name	Strength
CELLULOSE, MICRO CRYSTALLINE (UNII: OP1R32D61U)	
LACTOSE MONO HYDRATE (UNII: EWQ57Q8I5X)	
SILICON DIO XIDE (UNII: ETJ7Z6 XBU4)	
HYDROXYPROPYL CELLULOSE (70000 WAMW) (UNII: 66O7AQV0RT)	
PO VIDO NE (UNII: FZ989 GH94E)	
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	

Product Characteristics				
Color	WHITE (White to off-white)	Score	no score	
Shape	CAPSULE	Size	16 mm	
Flavor		Imprint Code	INCY;20	
Contains				

ı	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1 NE 60	OC:50881-020-	60 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	11/16/2011	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA202192	11/16/2011	

JAKAFI

ruxolitinib tablet

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:50881-025
Route of Administration	ORAL		

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
RUXOLITINIB (UNII: 82S8X8XX8H) (RUXOLITINIB - UNII:82S8X8XX8H)	RUXOLITINIB	25.0 mg		

Inactive Ingredients		
Ingredient Name	Strength	
CELLULO SE, MICRO CRYSTALLINE (UNII: OP1R32D61U)		
LACTOSE MONOHYDRATE (UNII: EWQ57Q8I5X)		
SILICON DIO XIDE (UNII: ETJ7Z6 XBU4)		
HYDROXYPROPYL CELLULOSE (70000 WAMW) (UNII: 6607AQV0RT)		
PO VIDO NE (UNII: FZ989 GH94E)		
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)		
MAGNESIUM STEARATE (UNII: 70097M6I30)		

Product Characteristics			
Color	WHITE (White to off-white)	Score	no score
Shape	OVAL	Size	18 mm
Flavor		Imprint Code	INCY;25
Contains			

ı	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:50881-025- 60	60 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	11/16/2011	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
NDA	NDA202192	11/16/2011		

Labeler - Incyte Corporation (556967347)

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